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




EDGE ARTICLE

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Realizing long range π -conjugation in phenanthrene and phenanthrene-based molecular crystals for anomalous piezoluminescence

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Realizing long range π -conjugation in phenanthrene and phenanthrene-based molecular crystals for anomalous piezoluminescence†

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Unlike the known aggregation-caused quenching (ACQ) that the enhancement of π - π interactions in rigid organic molecules usually decreases the luminescent emission, here we show that an intermolecular "head-to-head" π - π interaction in the phenanthrene crystal, forming the so-called "transannular effect", could result in a higher degree of electron delocalization and thus photoluminescent emission enhancement. Such a transannular effect is molecular configuration and stacking dependent, which is absent in the isomers of phenanthrene but can be realized again in the designed phenanthrene-based cocrystals. The transannular effect becomes more significant upon compression and causes anomalous piezoluminescent enhancement in the crystals. Our findings thus provide new insights into the effects of π - π interactions on luminescence emission and also offer new pathways for designing efficient aggregation-induced emission (AIE) materials to advance their applications.

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Introduction

Solid-state luminescent materials have been attracting intensive research interest because of their wide range of applications, such as optoelectronics,¹⁻⁴ data storage^{5,6} and theranostics.⁷⁻⁹ Aggregation-induced emission (AIE) has been considered a dominant mechanism for designing organic solid luminescent materials for many applications.¹⁰⁻¹⁹ In general, AIE molecules should be non-rigid with active intramolecular motions, and they usually show no emission in solutions at the molecular level but become brightly emissive in mesoscopic aggregated states.²⁰⁻²² For the AIE mechanism, it is generally accepted that the restriction of intramolecular motions (RIM) could reduce the energy dissipation channels of the non-radiative transition, enhance the radiative transition process and thus increase the emission in the aggregation state.^{14,18}

Compared to non-rigid organic luminescent molecules, rigid organic molecules could exhibit additional advantages such as higher thermal stability and structural stability,^{23,24} but rigid organic molecules usually show aggregation-caused quenching (ACQ)^{13,14} in the aggregated state while becoming highly emissive in the isolated form.²⁵ In this case, the π - π stacking enhancement that should affect the π electron behaviors during excitation and radiation processes is considered as the main

reason responsible for the ACQ.^{26,27} Therefore, how to develop a new strategy to realize solid-state luminescent materials with high efficiency for new functionalities (such as piezoluminescence) and extended practical applications by using rigid organic molecules is highly desirable but remains challenging.

Numerous studies show that the delocalization and flowability of π electrons in organic luminescent molecules play important roles in their optical properties.²⁸⁻³⁰ Graphene can be considered as an extremely large two-dimensional "conjugated molecule" constructed by substantial aromatic carbon rings,^{31,32} in which π electrons exhibit high delocalization and also high flowability, leading to excellent electrical properties³³⁻³⁵ but the absence of fluorescent properties. In contrast, when the number of aromatic rings decreases to the molecular level (such as polycyclic aromatic hydrocarbons, PAHs), π electrons tend to display heightened localization and reduced flowability because of the break of long range π -conjugation, which causes new properties and brings fluorescence emission.³⁶⁻³⁸ Therefore, a possible way to design solid luminescent materials using rigid organic molecules is to promote the delocalization of π electrons without increasing their flowability, which remains a challenging task but is a new strategy different from those by decreasing π - π stacking. However, this has not been explored for the design of efficient luminescent materials.

High-pressure can be used to precisely tune intermolecular interactions, altering molecular vibration and rotation modes and electronic excited states, and thus manipulating the emission behaviors.³⁹⁻⁴⁴ It is also an efficient technique to tune the delocalization of π electrons and amplify the related effects that

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are weak at ambient pressure. In this study, we discovered that the “transannular effect”,^{45–47} a type of weak π – π interaction, occurs in phenanthrene molecular crystals because of their unique molecular configuration. Such an effect becomes more significant upon compression and causes anomalous piezoluminescent enhancement in molecular crystals, which was also observed in phenanthrene-based cocrystals but absent in their isomers. The transannular effect could facilitate delocalization by generating long range π -conjugation in phenanthrene crystals while the flowability of π electrons is still limited within a single molecule, leading to an increase in radiative transition rates and emission intensity. Our finding not only adds another mechanism for aggregation-induced emission enhancement in luminescent materials but also presents an efficient strategy for the design of novel piezoluminescent materials.

Results and discussion

Experimental and theoretical details are presented in the ESI,[†] and structural information for all crystals and cocrystals involved in this work is presented in Fig. S1.[†] Fig. 1(a) displays the high-pressure photoluminescence (PL) spectra of phenanthrene crystals obtained by 360 nm excitation. The phenanthrene crystal shows split emission bands, similar to those of single molecules, which agrees well with the weak π – π stacking due to the bent geometry and a “herringbone” pattern⁴⁸ molecular arrangement in the phenanthrene crystal at ambient pressure (Fig. 1(f)). As shown in Fig. 1(b), phenanthrene shows anomalous emission enhancement at 1 atm–1.65 GPa as pressure increases, while at above 1.65 GPa, the emission exhibits the ACQ phenomenon (Fig. S2[†]). The quenched emission is

primarily attributed to the compression-induced strengthening of intermolecular interactions, which results in more electron internal conversion and less radiation. *In situ* ultraviolet-visible (UV-vis) absorption spectra demonstrate that the energy gap was narrowed upon compression (Fig. 1(d and e)), leading to the red shift of emission.

The pressure-dependent PL intensities of phenanthrene crystals upon compression are shown in Fig. 1(c), revealing a significant intensity enhancement of about 5-fold up to 1.65 GPa. To elucidate the underlying reasons for such an anomalous emission enhancement, *in situ* infrared (IR) and Raman spectra measurements were performed to investigate the intra- and intermolecular interactions in the phenanthrene crystal under high pressure. The experimental results match the calculated results well (Fig. S3(c) and (d)[†]), and as shown in Fig. 2(a) and S4,[†] all IR absorption peaks are blue-shifted upon compression, indicating an increase in the intramolecular vibrational energies of phenanthrene and an increase of non-radiative processes.

We noticed that the IR absorption intensity at approximately 793 cm^{-1} , which represents the carbon skeleton out-of-plane bending vibration of the phenanthrene molecule according to our calculation, increases with pressure. The corresponding intensities of the 793 cm^{-1} mode as a function of pressure are plotted and shown in Fig. 2(b). This vibration mode exhibits a high degree of symmetry, and its intensity enhancement upon compression indicates that the molecule's dipole moment changes. This change may be due to the increased electronic delocalization of the intermediate aromatic ring because of the non-linear molecular structure of phenanthrene. Our Raman measurements show that all Raman modes are blue-shifted

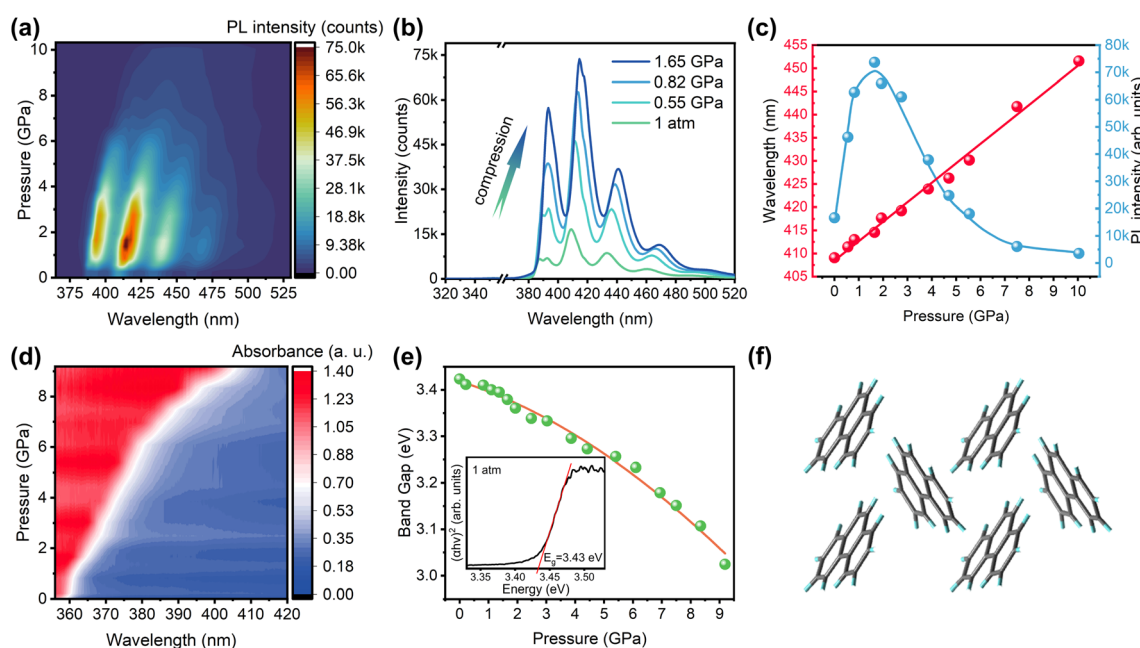


Fig. 1 (a) 2D colormap of pressure-dependent PL intensity and wavelength of the phenanthrene crystal. (b) *In situ* PL spectra of the phenanthrene crystal below 1.65 GPa. (c) Pressure-dependent PL wavelength and intensities. (d) 2D colormap of high-pressure UV-vis absorption spectra of the phenanthrene crystal. (e) Schematic illustrations of the band gap changes. (f) The sketch map for molecular packing of the phenanthrene crystal.



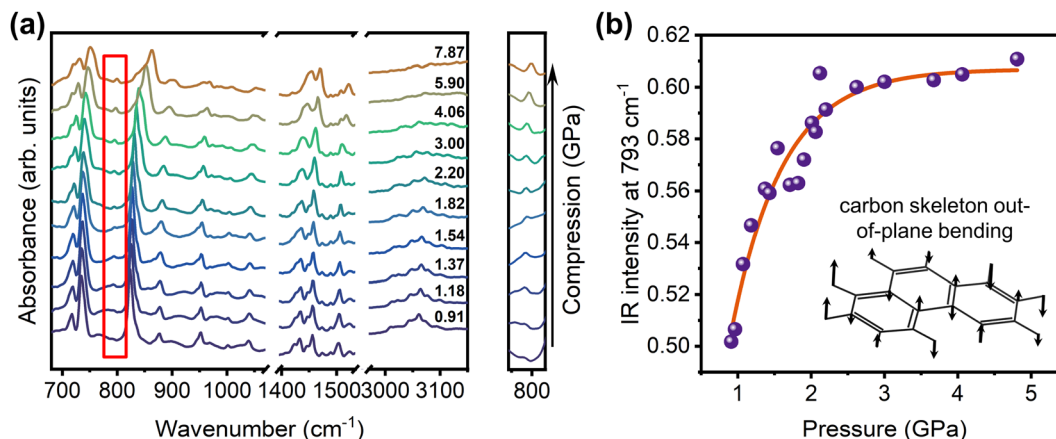


Fig. 2 (a) *In situ* IR spectra of the phenanthrene crystal upon compression. (b) The pressure-dependent IR absorption intensities around 793 cm^{-1} and the schematic diagram of the corresponding vibrational mode. The relevant vibrations are marked in the diagrams (a).

upon compression, which is consistent with our IR results (Fig. S3[†]). This clearly suggests that the general RIM mechanism cannot explain why phenanthrene exhibits such anomalous emission upon compression. Note that the Raman peaks around 100 cm^{-1} (see the marked regions in Fig. S3[†]) from out-of-plane carbon skeleton deformation vibrations exhibit a rapid blue shift at low pressure, implying significant interactions between the neighboring phenanthrene molecules, which may lead to a change in the π -electronic distribution on phenanthrene molecules. Combining our results from IR and Raman measurements, we suggest there is an active intermolecular

nonbonding interaction – transannular effect. Note that previous studies show that the transannular effect can indeed alter the electronic structures of the aromatic system.^{45,46,49,50} In our case, such a unique interaction that most likely occurs between phenanthrene molecules could lead to greater delocalization of π electrons, which can increase the radiative transition rates and result in an anomalous piezoluminescent enhancement.

To understand if the molecular stacking could favor the transannular effect in the crystal upon compression, a high-pressure X-ray diffraction (XRD) experiment has been carried

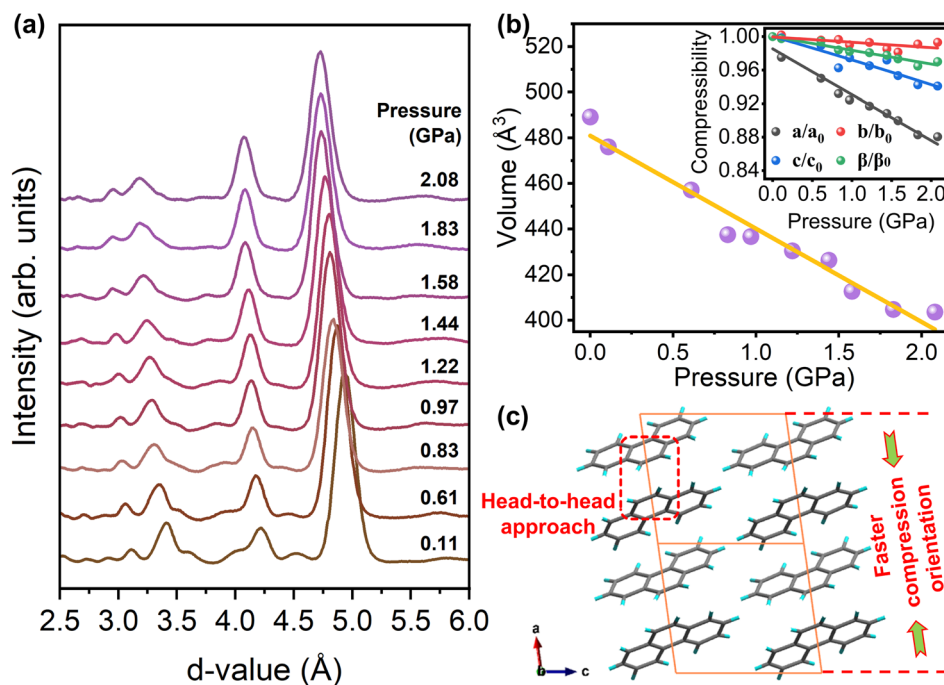


Fig. 3 (a) High-pressure XRD patterns of the phenanthrene crystal below 2.08 GPa. (b) Plotted curves for the unit cell volume of phenanthrene as a function of pressure. Inset shows the compression rate of lattice constants as pressure increases, which is given by the monoclinic $P2_1$ structure. (c) The crystal structure viewed along the b -axis, in which the marked regions represent the head-to-head approach between molecules. The XRD patterns are analyzed by JADE.



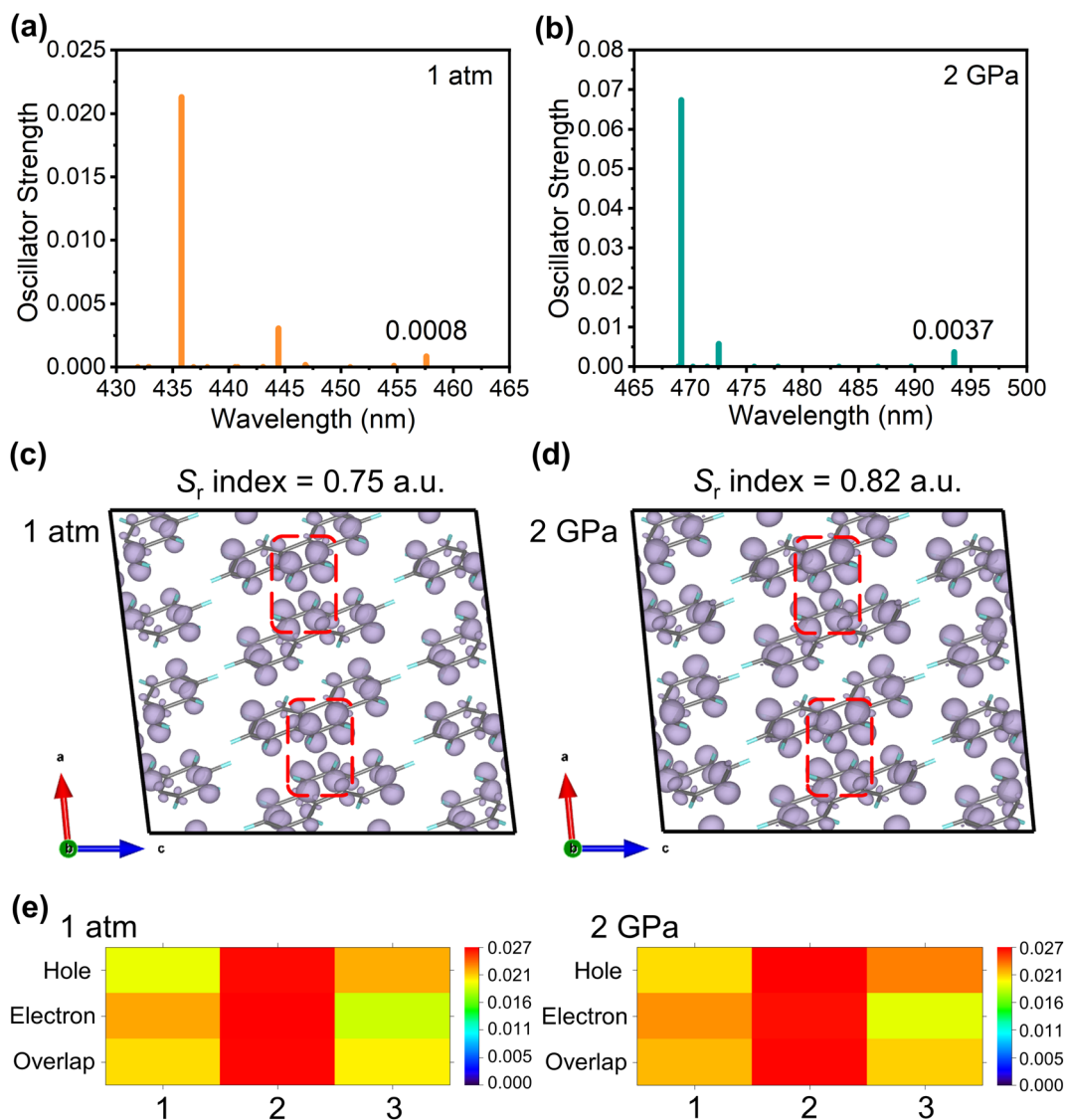


Fig. 4 (a and b) The emission oscillator strengths of phenanthrene calculated at 1 atm and 2 GPa. (c and d) The overlap function $S_r(r)$ during the $S_1 \rightarrow S_0$ transition at 1 atm and 2 GPa (isosurface level = 0.00016 a.u.), the marked regions represent the head-to-head areas between molecules. (e) The contributions of the three parts of the phenanthrene molecule (rings 1, 2, 3) to the hole, electron, and orbital overlap during the $S_1 \rightarrow S_0$ transition at 1 atm and 2 GPa.

comparison, the linear isomer of phenanthrene, anthracene, was studied and it exhibits a normal ACQ behavior accompanied by red-shifted emission upon compression in both experiments and calculations (Fig. S7(b-d)[†]), despite anthracene and phenanthrene having very similar molecular packing in the corresponding crystals (Fig. S7(a)[†]). The molecular configuration-dependent emission responses can be understood by the fact that the linear anthracene can easily form an excimer^{60,61} and thus reduce the emission under high pressure, while the bent geometry is more favorable to suppress the increased π - π stacking between parallel phenanthrene molecules upon compression and provides a stronger steric effect. These differences in molecular configuration may be the reason why transannular effects do not occur in anthracene crystals.

We further demonstrate that the transannular effect could be either preserved or suppressed by introducing another

component to form piezoluminescent cocrystals. Considering from molecular packing, we insert 1,2,4,5-tetracyanobenzene (TCNB) into phenanthrene (Phe) crystals, forming Phe-TCNB cocrystals. The TCNB could serve as an acceptor and hold the cocrystal's lowest unoccupied molecular orbital (LUMO), thereby simplifying the energy transfer route (charge-transfer (CT) emission). The TCNB could also prevent the face-to-face packing of phenanthrene molecules and thus leave only their head-to-head interactions (Fig. 5(a)). Our high-pressure PL spectra (Fig. 5(b)) reveal that the Phe-TCNB cocrystal indeed exhibits emission enhancement at a pressure range of 1 atm–1.77 GPa, but no structural phase change occurs in the cocrystals (Fig. S8[†]). The high-pressure IR absorption spectra of the Phe-TCNB cocrystal (Fig. S9[†]) also show that all IR absorption peaks were blue-shifted as pressure increased, indicating that the nonradiative energy dissipation of the molecule is not suppressed. Instead,



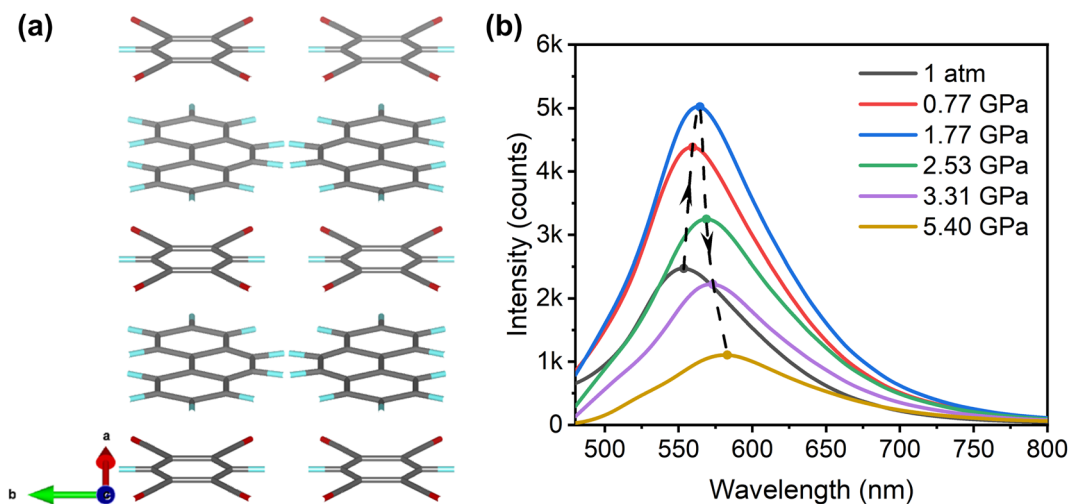


Fig. 5 (a) The sketch map for molecular packing of Phe-TCNB. (b) High-pressure PL spectra of Phe-TCNB.

the 796 cm^{-1} and 1602 cm^{-1} absorption peaks of Phe-TCNB from the carbon skeleton out-of-plane and in-plane bending display anomalous intensity enhancement as pressure increases, giving a strong indication for the transannular effect in this cocrystal. Note that when phenanthrene was replaced by its linear isomer anthracene (Ant), forming the Ant-TCNB cocrystal, emission quenching was observed upon compression (Fig. S10†). We could also destroy the head-to-head interactions between the phenanthrene molecules by inserting anthracene in Phe-TCNB cocrystals, forming a ternary $\text{Ant}_{0.5}\text{-Phe}_{0.5}\text{-TCNB}$ cocrystal. In this case, the transannular effect was suppressed and emission quenching was observed upon compression (Fig. S11†). These results thus demonstrate that the transannular effect can be applicable to different luminescent materials, offering new pathways for designing more efficient organic luminescent materials for different applications.

Experimental and computational details

Materials source, synthesis and crystal structure

Phenanthrene (Phe, 98%) was purchased from Alfa Aesar, anthracene (Ant, 99%) was purchased from MACKLIN, and 1,2,4,5-tetracyanobenzene (TCNB, 97%) was purchased from Tokyo Chemical Industry Co., Ltd (TCI). All of the chemicals were used as received without further purification. The Phe-TCNB, Ant-TCNB and $\text{Ant}_{0.5}\text{-Phe}_{0.5}\text{-TCNB}$ cocrystals were prepared by solvent evaporation methods. Identical molar masses of phenanthrene and TCNB were dissolved in excess tetrahydrofuran (THF) solutions and ultrasonicated for 12 minutes. The yellow stick-like transparent sample Phe-TCNB was obtained after evaporation of solvent from the solution after 3–4 days under ambient conditions. Ant-TCNB and $\text{Ant}_{0.5}\text{-Phe}_{0.5}\text{-TCNB}$ cocrystals can be obtained by the same method. The initial crystal structures of our phenanthrene and anthracene were obtained from the Cambridge Crystallographic Data Centre (CCDC, no. 1232373 and no. 950158).

In situ high-pressure experiments

High-pressure experiments were performed in a diamond anvil cell (DAC). Samples were loaded into a $120\text{ }\mu\text{m}$ diameter hole drilled in the T301 stainless-steel gasket. Pressure was calibrated by the fluorescence emission of ruby in the sample chamber.⁶² The pressure transmitting medium (PTM) used in the high-pressure PL, XRD, Raman and UV-vis absorption measurements was silicone oil, while it was KBr in the high-pressure IR measurements. A home-built integrated optical measurement system was used to collect high-pressure PL spectra and ultraviolet-visible (UV-vis) absorption spectra of phenanthrene and anthracene, and a semiconductor UV laser and a deuterium-halogen lamp were used as the excitation sources for the PL and absorption spectra, respectively, along with a Horiba Jobin Yvon iHR320 spectrometer; the PL excitation laser had a wavelength of 360 nm . And PL measurements of Phe-TCNB, Ant-TCNB and $\text{Ant}_{0.5}\text{-Phe}_{0.5}\text{-TCNB}$ were performed on a Raman spectrometer equipped with a CCD detector (Renishaw in Via) in fluorescence mode, and a 514 nm line of a Cobolt Fandango™ laser was used for the excitation source. Infrared measurements were carried out using a Bruker Vertex 80 V spectrometer with a liquid nitrogen-cooled MCT detector. *In situ* high-pressure X-ray diffraction measurements were performed using a Rigaku Synergy Custom FR-X ($\lambda = 0.7093\text{ \AA}$), while ambient-pressure X-ray diffraction measurements were performed using a Rigaku MicroMax-007 HF ($\lambda = 1.5418\text{ \AA}$). High-pressure Raman spectra were collected using a LabRAM HR Evolution spectrometer (HORIBA Jobin-Yvon) excited by a 473 nm laser.

Computational details

The vibrational analysis was performed using the CASTEP⁶³ module in the Materials Studio package. The exchange and correlation of electrons were treated by the generalized gradient approximation (GGA) with the Perdew–Burke–Ernzerhof (PBE) functional, and the OTFG norm-conserving pseudo-potentials were used for calculations. We used the CP2K software



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